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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/787,368	02/26/2004	Hyun-Woo Kim	8028-42 (SPX200304-0017US	7951	
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F. CHAU & A	ASSOCIATES, LLC		WALKE, AMANDA C		
WOODBURY, NY 11797			ART UNIT	PAPER NUMBER	
			1752		
			DATE MAILED: 05/16/2006		

Please find below and/or attached an Office communication concerning this application or proceeding.

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		Application No.	Applicant(s)	μ=		
		10/787,368	KIM			
	Office Action Summary	Examiner	Art Unit			
		Amanda C. Walke	1752			
Period fo	The MAILING DATE of this communication app or Reply	pears on the cover sheet with the	correspondence addre	ess		
WHIC - Exte after - If NC - Failu Any	ORTENED STATUTORY PERIOD FOR REPLY CHEVER IS LONGER, FROM THE MAILING DATE of time may be available under the provisions of 37 CFR 1.1: SIX (6) MONTHS from the mailing date of this communication. Opened for reply is specified above, the maximum statutory period were to reply within the set or extended period for reply will, by statute reply received by the Office later than three months after the mailing ed patent term adjustment. See 37 CFR 1.704(b).	ATE OF THIS COMMUNICATIO 36(a). In no event, however, may a reply be ti will apply and will expire SIX (6) MONTHS fror , cause the application to become ABANDON	N. imely filed in the mailing date of this comm ED (35 U.S.C. § 133).			
Status						
1)⊠	Responsive to communication(s) filed on <u>09 M</u>	larch 2006.				
2a)⊠	This action is FINAL . 2b) This	action is non-final.				
3)	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is					
	closed in accordance with the practice under E	Ex parte Quayle, 1935 C.D. 11, 4	53 O.G. 213.			
Dispositi	ion of Claims					
5)□ 6)⊠ 7)□	Claim(s) 1,3-5,10-16,21-33 and 35-37 is/are per 4a) Of the above claim(s) is/are withdraw Claim(s) is/are allowed. Claim(s) 1,3-5,10-16,21-33 and 35-37 is/are reclaim(s) is/are objected to. Claim(s) are subject to restriction and/o	wn from consideration.				
Applicati	ion Papers					
9) <u> </u> 10) <u> </u>	The specification is objected to by the Examine The drawing(s) filed on is/are: a) accomplicant may not request that any objection to the Replacement drawing sheet(s) including the correct The oath or declaration is objected to by the Example.	epted or b) objected to by the drawing(s) be held in abeyance. So tion is required if the drawing(s) is of	ee 37 CFR 1.85(a). bjected to. See 37 CFR			
Priority ι	under 35 U.S.C. § 119					
12)⊠ a)i	Acknowledgment is made of a claim for foreign All b) Some * c) None of: 1. Certified copies of the priority document: 2. Certified copies of the priority document: 3. Copies of the certified copies of the priority document: application from the International Bureau See the attached detailed Office action for a list	s have been received. s have been received in Applica rity documents have been receiv u (PCT Rule 17.2(a)).	tion No ved in this National Sta	age		
	e of References Cited (PTO-892)	4) Interview Summar				
3) 🔲 Infor	te of Draftsperson's Patent Drawing Review (PTO-948) mation Disclosure Statement(s) (PTO-1449 or PTO/SB/08) or No(s)/Mail Date	Paper No(s)/Mail I 5) Notice of Informal 6) Other:	Date Patent Application (PTO-1	52)		

DETAILED ACTION

Claim Rejections - 35 USC § 103

- 1. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- 2. Claims 1, 3-5, 11-16, 22-33, and 35-37 are rejected under 35 U.S.C. 103(a) as being unpatentable over Foster et al (6,054,248) in view of Gruber et al (4,276,136), Ryang et al (6,261,687), or Tamura et al (2004/0009428) in view of Schaedeli et al (6,146,793).

Foster et al teaches a thermally curable polymer composition comprising a hydroxylcontaining polymer and a polyfunctional isocyanate as a crosslinking agent, which is particularly
useful in photolithographic coating of substrates. The thermally curable polymer composition
may be dissolved in a solvent and used as an undercoat layer in deep UV lithography.

Furthermore, the present invention further relates to a process for using the photolithographic
coated substrate for the production of relief structures.

The reference teaches that "The thermally curable polymer composition also comprises a hydroxyl-containing polymer. Suitable examples of hydroxyl-containing polymers are polymers comprising monomer units selected from the group consisting of: cyclohexanol, hydroxyalkyl acrylate or methacrylate, hydroxycycloalkyl acrylate or methacrylate, hydroxycycloalkyl acrylate or methacrylate, allyl alcohol and the like. This invention also contemplates copolymers, terpolymers, etc. of the foregoing named polymers.

Preferably, polymers comprising monomer units of cyclohexanol, hydroxyalkyl acrylate or methacrylate, and hydroxycycloalkyl acrylate or methacrylate have a number average molecular weight of about 9000 to 38,000, more preferably about 14,000 to 30,000 and most preferably about 18,000 to 22,000.

In addition, the thermally curable polymer composition may also further comprise monomer units of cycloaliphatic esters of acrylic or methacrylic acid. Suitable examples of monomer units of cycloaliphatic esters of acrylic or methacrylic acid are cyclohexyl acrylate or methacrylate, 4-tert-butylcyclohexyl acrylate or methacrylate, isobomyl acrylate or methacrylate and adamantyl acrylate or methacrylate and the like. The preferred monomer units of cycloaliphatic ester of acrylic or methacrylic acid are isobornyl acrylate or methacrylate.

Furthermore, the hydroxyl-containing polymer may further comprise aromatic monomer units, preferably styrene.

In a preferred embodiment, the thermally curable polymer composition comprises monomer units of (a) a cycloaliphatic ester of acrylic or methacrylic acid; (b) a sole aliphatic ester of acrylic or methacrylic acid consisting of hydroxyalkyl acrylate or methacrylate; and (c) a polyfunctional isocyanate, wherein the polymer has a number average molecular weight of about 9000 to 38,000, preferably about 14,000 to 30,000, more preferably about 18,000 to 22,000.

Other preferred copolymers useful in the thermally curable polymer composition are a copolymer of styrene and allyl alcohol with a weight average molecular weight of about 2000 to 20,000, preferably 2,000 to 10000; and a terpolymer of hydroxyalkyl acrylate or methacrylate, cycloaliphatic esters of acrylic or methacrylic acid and styrene with a number average molecular

weights of about 9000 to 38,000, preferably about 14,000 to 30,000, more preferably about 18,000 to 22,000.

Examples of suitable hydroxyalkyl acrylate or methacrylates are hydroxymethyl acrylate or methacrylate, 2-hydroxyethyl acrylate or methacrylate, 3-hydroxypropyl acrylate or methacrylate, 4-hydroxybutyl acrylate or methacrylate, 5-hydroxypentyl acrylate or methacrylate, and 6-hydroxyhexyl acrylate or methacrylate and the like. Preferably, the hydroxyalkyl acrylate or methacrylate contains primary hydroxyl groups, although secondary alcohol groups or mixtures of primary and secondary alcohol groups may be used. Suitable examples of secondary alcohols are 2-hydroxy-2-methylethyl acrylate or methacrylate, 3-hydroxy-3-methylpropyl acrylate, 4-hydroxy-4-methylbutyl acrylate or methacrylate, 5-hydroxy-5-methyl propyl acrylate or methacrylate, and the like. The preferred hydroxyalkyl acrylate or methacrylate is 2-hydroxyethyl acrylate or methacrylate.

The thermally curable polymer composition of the present invention should not begin significant cross-linking until it reaches a temperature of about 50.degree. C. Significant cross-linking below 50.degree. C. may lead to gel formation at room temperature, which will reduce the composition's shelf life. Gel formation results in non-uniform coatings and linewidth variations across the substrate when the thermally curable polymer composition is used as an undercoat layer in microlithography.

Polymer 1 comprises about 60 to 80 mole % of isobomyl acrylate or methacrylate monomer units and about 20 to 40 mole % of 2-hydroxyethyl acrylate or methacrylate monomer units.

Polymer 2 comprises about 30 to 40 mole % of isobomyl acrylate or methacrylate monomer units, about 10 to 30 mole % of 2-hydroxyethyl acrylate or methacrylate monomer units, and about 30 to 50 mole % of styrene monomer units.

Polymer 3 comprises about 39-60 mole % of styrene monomer units and about 40 to 61 mole % of allyl alcohol monomer units.

The polymerization of the polymers described above may be carried out by any suitable polymerization process such as free radical polymerization. The number average molecular weight of the Polymers 1 and 2 are about 9,000 to 38,000, preferably about 14,000 to 20,000, and more preferable about 18,000 to 22,000.

The present invention also relates to a photolithographic coated substrate comprising: a substrate, a thermally cured undercoat composition on the substrate, and a radiation-sensitive resist topcoat on the thermally cured undercoat composition.

The thermally cured undercoat composition comprises the thermally curable polymer composition comprising a hydroxyl-containing polymer and a polyfunctional isocyanate cross-linking agent that has been heated to form a cross-linked matrix. Any of the polymers described above may be used as the hydroxyl-containing polymer. Preferably, the hydroxyl-containing polymer is selected from Polymers 1, 2 or 3.

The present invention further relates to a process for using the photolithographic coated substrate for the production of relief structures comprising the steps of: providing the photolithographic coated substrate, imagewise exposing the radiation-sensitive resist topcoat to actinic radiation; and forming a resist image by developing the radiation-sensitive resist topcoat with a developer to form open areas in the radiation-sensitive resist topcoat. In addition, the

thermally cured undercoat composition may be removed in the open areas of the developed radiation-sensitive resist topcoat by any suitable process such as oxygen plasma etching to form an image in the thermally cured undercoat composition.

One advantage of the thermally curable polymer composition is that it may be cured at a temperature of less than about 250 degrees C and for a time less than about 180 seconds. This make it particularly useful as an undercoat layer for a resist system where temperature and time constraints are important for commercial viability.

Both the undercoat and the radiation-sensitive compositions are uniformly applied to a substrate by known coating methods. The compositions are solubilized in an organic solvent and the coatings may be applied by spin-coating, dipping, knife coating, lamination, brushing, spraying, and reverse-roll coating. The coating thickness range generally covers values of about 0.1 to more than 10 microns, and more preferably from about 0.1 to 1.5 microns for the radiation-sensitive resist and about 0.3 to 3.0 microns for the undercoat layer. After the coating operation, the solvent is generally removed by curing or drying.

Suitable solvents for both the undercoat and topcoat radiation-sensitive compositions include ketones, ethers and esters, such as methyl ethyl ketone, methyl isobutyl ketone, 2-heptanone, cyclopentanone, cyclopentanone, 2-methoxy-1-propylene acetate, 2-methoxyethanol, 2-ethoxyethanol, 2-ethoxyethyl acetate, 1-methoxy-2-propyl acetate, 1,2-dimethoxy ethane ethyl acetate, cellosolve acetate, propylene glycol monoethyl ether acetate, propylene glycol methyl ether acetate, methyl lactate, ethyl lactate, methyl pyruvate, ethyl pyruvate, methyl 3-methoxypropionate, ethyl 3-methoxypropionate, N-methyl-2-pyrrolidone, 1,4-dioxane, ethylene

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glycol monoisopropyl ether, diethylene glycol monoethyl ether, diethylene glycol monomethyl ether, diethylene glycol dimethyl ether, and the like.

The radiation-sensitive resist topcoat of the present invention may be any suitable radiation-sensitive resist. It is typically a chemically amplified resist sensitive to radiation in the deep UV region.

The radiation-sensitive resist will also contain a photoacid generating (PAG) compound. The PAG compounds may be of any suitable type such as sulfonium or iodonium salts, nitrobenzyl esters, imidosulfonates esters and the like. Typically the PAG will be in an amount of about 1 to 10% based on the weight of the polymer.

For the production of relief structures, the radiation-sensitive resist is imagewise exposed to actinic radiation. The term 'imagewise' exposure includes both exposure through a photomask containing a predetermined pattern, exposure by means of a computer controlled laser beam which is moved over the surface of the coated substrate, exposure by means of computer-controlled electron beams, and exposure by means of X-rays or UV rays through a corresponding mask. The imagewise exposure generates acid in the exposed regions of the resist which cleaves the acid labile groups resulting in a polymer which is aqueous soluble. Typically, after imagewise exposure, the chemically amplified resist will be subjected to a post exposure heating treatment that virtually completes the reaction of the photoacid generator with the acid labile groups.

After imagewise exposure and any heat treatment of the material, the exposed areas of the top radiation-sensitive resist are typically removed by dissolution in a aqueous developer.

The choice of the particular developer depends on the type of photoresist; in particular on the

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nature of the polymer resin or the photolysis products generated. The developer can comprise aqueous solutions of bases to which organic solvents or mixtures thereof may have been added. Particularly preferred developers are aqueous alkaline solutions. These include, for example, aqueous solutions of alkali metal silicates, phosphates, hydroxides and carbonates, but in particular of tetra alkylammonium hydroxides, and more preferably tetramethylammonium hydroxide (TMAH). If desired, relatively small amounts of wetting agents and/or organic solvents can also be added to these solutions.

The radiation-sensitive resist used for the bilayer process described above will typically contain silicon or have silicon incorporated into the resist after development. After images are formed in the radiation-sensitive resist, the substrate will be placed in an plasma-etching environment comprising oxygen so that the underlayer coating will be removed. The silicon incorporated in the radiation-sensitive resist forms silicon dioxide when exposed to an oxygen plasma and protects it from being etched so that relief structures can be formed in the undercoat layer.

After the oxygen plasma step, the substrate carrying the bilayer relief structure is generally subjected to at least one further treatment step which changes the substrate in areas not covered by the bilayer coating. Typically, this can be implantation of a dopant, deposition of another material on the substrate or an etching of the substrate. This is usually followed by the removal of the resist coating from the substrate typically by a fluorine/oxygen plasma etch. (column 4, line 1 to column 8, line 16).

The reference teaches that the underlayer and resist layer may be baked or cured after coating by any known method. But the reference fails to specifically teach ebeam curing.

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Gruber et al, Ryang et al, and Tamura et al all teach that similar polymeric curable compositions are cured by ebeam.

Given the teachings of the reference that clearly disclose polymers meeting the instant claim limitations employed in an undercoat for a bilayer resist and a method of forming a pattern using the same as described by the instant invention, it would have been obvious to one of ordinary skill in the art to prepare the material of Foster et al choosing to employ ebeam to cure the thermally curable underlayer given that it is taught to be conventional by Tamura et al, Ryang et al, and Gruber et al.

Foster et al has been discussed above, and while the reference clearly anticipates the use of the methacrylate polymer with an aromatice polymer (styrene), it fails to teach the use of a novolac resin.

Schaedeli et al disclose a bilayer system for use in 193 nm photolithography. The invention also relates to radiation sensitive photoresist compositions comprising a new terpolymer and to a process for the lithographic treatment of a substrate by means of the photoresist composition, as well as a process for the production of electronic components using the new terpolymer in a radiation sensitive photoresist top layer coating in a 193 nm lithographic system.

The reference further teaches that "any suitable film-forming organic polymers can be used as the film-forming organic material for the first coating (undercoat layer) with the use of the multilayer technique. Particularly preferred are phenolic resins, particularly novolak resins, such as formaldehyde cresol or formaldehyde phenol novolaks, polyimide resins, poly(meth) acrylate resins and styrene-allyl alcohol copolymer resins, the latter being preferred. "

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Given the teachings of the reference, it would have been obvious to one of ordinary skill in the art to prepare the material of Foster et al choosing to employ any phenolic resin in the undercoat layer such as a novolak resin given the teaching of Schaedeli et al that it is conventional, with reasonable expectation of achieving a material having high resolution.

5. Claims 10 and 21 are rejected under 35 U.S.C. 103(a) as being unpatentable over Foster et al in view of Ryang et al, Gruber et al, or Tamura et al in view of Wong et al (6,319,655).

Foster et al has been discussed above, and while the reference teaches that the material is suitable for exposure at the 193 nm region, the reference fails to teach or suggest suitable exposure doses.

Wong et al disclose a resist that typical UV exposure doses fall within the limitations of the instant claims, as do the typical electron beam exposures (column 7, line 45 to column 8, line 64).

Given the teachings of the reference it would have been obvious to one of ordinary skill in the art to prepare the material of Foster et al using the exposure doses taught by Wong et al which are conventional doses, with reasonable expectation of achieving a material having high resolution.

Response to Arguments

3. Applicant's arguments filed 3/9/2006 have been fully considered but they are not persuasive. Applicant has argued that the amendments to the claims overcome the 102 rejection (which has been dropped), but that any 103 rejection would not be appropriate as there is no motivation to combine the references of record. As discussed above, the Foster reference does

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teach a layer having polymers (see specific polymers cited above which clearly have various (meth)acrylates with styrene)that are instantly disclosed and teach that other known polymers may be added, and Schaedeli et al teaches that a various novolac resins are known and preferably added to such compositions having styrene and (meth)acylate polymers. The Gruber, Ryang, and Tamura, as atuaght above, Gruber et al, Ryang et al, and Tamura et al all teach that similar polymeric curable compositions are cured by ebeam. These references clearly disclose polymers meeting the instant claim limitations employed in an undercoat for a bilayer resist and a method of forming a pattern using the same as described by the instant invention, thus it would have been obvious to one of ordinary skill in the art to expose to ebeam to cute the underlayer, and the rejection is maintained.

Conclusion

3. Applicant's amendment necessitated the new ground(s) of rejection presented in this Office action. Accordingly, **THIS ACTION IS MADE FINAL**. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the date of this final action.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Amanda C. Walke whose telephone number is 571-272-1337. The examiner can normally be reached on M-R 5:30-4.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Cynthia Kelly can be reached on 571-272-1526. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free).

Ay Whom Wilk Amanda C Walke Primary Examiner Art Unit 1752

ACW May 9, 2006